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Office européen des brevets

1) Publication number:

0 468 520 A3

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EUROPEAN PATENT APPLICATION

Application number: 91112601.9

1) Int. Cl. 4 A61K 31/70

22 Date of filing: 26.07.91

® Priority: 27.07.90 JP 197778/90

Date of publication of application: 29.01.92 Bulletin 92/05

Designated Contracting States:
DE FR GB IT

Date of deferred publication of the search report: 01.07.92 Bulletin 92/27

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Immunostimulatory remedies containing palindromic DNA sequences.

(7) Single-stranded, linear polydeoxyribonucleotides with a base number of 10 to 100 containing at least one structure represented by the following general formula:

 $5'-X_n \cdot \cdot \cdot X_3 X_2 X_1 Y_1 Y_2 Y_3 \cdot \cdot \cdot Y_n - 3'$ (1)

(wherein n is an integer from 3 to 50: X_1 , X_2 , X_3 , ..., X_n and Y_1 , Y_2 , Y_3 , ..., Y_n are each a monodeoxyribonucleotide; X_1 , X_2 , X_3 , ... and X_n may be the same or different nucleotides; and bases in X_1 and Y_1 , in X_2 and Y_2 , in X_3 and Y_3 , in ..., and in X_n and Y_n are complementary with each other as defined by Watson & Crick), and double-stranded, linear polydeoxyribonucleotides, in which at least one single-stranded, linear polydeoxyribonucleotide contains at least one structure represented by the general formula (I), both show strong im-

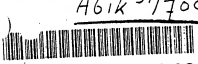
munostimulatory activity.

Thus, remedies containing, as active ingredient, such a specific polydeoxyribonuleotide as disecribed

above, or a salt thereof, are efficacious against malignant tumors, infectious diseases, immunodeficiency diseases and autoimmune diseases, with minimized side-effects.

EP 0 468 520 A3





(1) Publication number:

0 468 520 A2

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(1) Application number: 91112601.9

(1) Int. Cl.5: A61K 31/70

Date of filing: 26.07.91

3 Priority: 27.07.90 JP 197778/90

43 Date of publication of application: 29.01.92 Bulletin 92/05

Designated Contracting States: DE FR GB IT

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(S) Immunostimulatory remedies containing palindromic DNA sequences.

 Single-stranded, linear polydeoxyribonucleotides with a base number of 10 to 100 containing at least one structure represented by the following general formula:

5'-X_n···X₃X₂X₁Y₁Y₂Y₃···Y_n-3'

(wherein n is an integer from 3 to 50; X_1 , X_2 , X_3 , \cdots , X_n and Y_1 , Y_2 , Y_3 , \cdots , Y_n are each a monodeoxyribonucleotide; X_1 , X_2 , X_3 , \cdots and X_n may be the same or different nucleotides; and bases in X_1 and Y_1 , in X_2 and Y_2 , in X_3 and Y_3 , in \cdots , and in X_n and Y_n are complementary with each other as defined by Watson & Crick), and double-stranded, linear polydeoxyribonucleotides, in which at least one single-stranded. linear polydeoxyribonucleotide contains at least one structure represented by the general formula (I), both show strong immunostimulatory activity.

Thus, remedies containing, as active ingredient, such a specific polydeoxyribonuleotide as described above. or a salt thereof, are efficacious against malignant tumors, infectious diseases, immunodeficiency diseases and autoimmune diseases, with minimized side-effects.

The structure represented by the general formula (I) of this invention is a sequence of monodeoxyribonucleotides called palindromic structure, in which X₁ and Y₁, X₂ and Y₂, X₃ and Y₃, ***, and X_n and Y_n are complementary with each other as defined by Watson & Crick. A palindromic structure generally means one of the symmetric structures found in a double-stranded DNA, and these structures are the recognition site for many kinds of restriction enzymes. In this invention, this structural nomenclature commonly employed for double-stranded DNAs is used also for single-stranded DNAs, for convenience, and the structure represented by the general formula (I) is hereinafter called the palindromic structure or

palindromes.

What is to be noticed here is that the single- or double-stranded DNAs which are entirely composed of
What is to be noticed here is that the single- or double-stranded DNAs which are entirely composed of
alternately repeated sequence of only two types of monodeoxynucleotides (for example, G and C, or A and
T) cannot achieve the purpose of this invention. "n" in the general formula (I) is an integer from 3 to 50.

The following sequences are examples of the desirable structure of the present invention, wherein G is a deoxyguanylic acid, A is a deoxyadenylic acid, C is a deoxycytidylic acid and T is a deoxythymidylic acid, and wherein the left side is 5'-terminal and the right side is 3'-terminal in each sequence:

GAGCTC, TAGCTA, AGGCCT, CATATG, TGTACA, GTTAAC, GATATC,

GGGCCC, TTGCAA (n = 3)

GTAGCTAC, AAGGCCTT, GGATATCC, CAGGCCTG, GCATATGC, GTGTACAC,

AGTTAACT (n = 4)

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25 AGTAGCTACT, GAAGGCCTTC, AGGATATCCT, GCAGGCCTGC,

AGCATATGCT (n = 5)

To achieve the purpose of this invention, it is more desirable that one or more of the 5'-CG-3' structure be included in the structure represented by the general formula (I). As examples of such a structure, may be mentioned the following sequences:

CGATCG, ATCGAT, TCGCGA, AACGTT, GCGCGC, CGTACG, AGCGCT, CGGCCG, GACGTC, GTCGAC, CGCGCG, ACGCGT, CACGTG (n = 3

ACGATCGT, GATCGATC, ATCGCGAT, CAACGTTG, AGCGCGCT, ACGTACGT, TAGCGCTA, ACGGCCGT, CGACGTCG, CGTCGACG (n=4)

GACGATCGTC, CGATCGATCG, GATCGCGATC, GCAACGTTGC,

CAGCGCGCTG, GACGTACGTC, CTAGCGCTAG, GACGGCCGTC, ACGACGTCGT,

ACGTCGACGT, ACAACGTTGT (n = 5)

The single-stranded, linear DNA of this invention is an unbranched DNA molecule in which each of the compon nt monodeoxyribonucleotides is linked to the adjacent monodeoxyribonycleotide through a[5'-3'] phosphodiest r bond. The single-stranded, linear DNAs carrying less than six bases, which fail to satisfy the phosphodiest r bond. The single-stranded, linear DNAs carrying less than six bases, which fail to satisfy the general formula (I), are not satisfactory. As may be apparent from the Examples described later, the longer the chain length of single-stranded, linear DNA, the better will be their sult. However, the purpose of this invention may be sufficiently achieved with a DNA with a base number in the range from 10 to 100, because

functions of immune system are suppressed or lost, such as agammaglobulinemia and acquired immunodeficiency syndromes. Among the patients of these diseases, the morbidity of infectious diseases and malignant tumors is high, thus adversely affecting recuperation. DNAs of this invention, which are efficacious against malignant tumors and are also capable of inducing interferon, are expected to encourage the recuperation of the patients suffering immunodeficiency diseases by curing the malignant tumors and infectious diseases which are likely to concur in these patients.

Single- and double-stranded, linear DNAs of this invention may be administerd to animal and human bodies subcutaneously, intravenously, intramuscularly, intratumorally, orally or into the rectum, and the suitable administration route should be selected case by case depending on the type of disease and the conditions of the patient. For example, intratumoral or subcutaneous administration is preferable in the case of malignant tumors. The proper dose to humans is I to I000 mg/day when administered into the rectum or orally, and 0.01 to 100 mg/day when administered subcutaneously, intravenously, intratumorally or intramuscularly. Administration should be repeated once or twice per one to seven days, preferably once per one or two days, and the frequency of administration may be varied and the period of administration may

When administering single- or double-stranded, linear DNAs of this invention to animal and human be further prolonged, as required. bodies subcutaneously, intravenously, intramuscularly or intratumorally, it is preferable to appply it in the form of an injection prepared by dissolving the DNA in an aqueous solution which is nearly neutral (pH 5 to 8) with a physiological osmotic pressure. As examples of such an aqueous solution, may be mentioned the isotonic sodium chloride solution specified in Pharmacopoeia of Japan, and aqueous solutions containing salts, compounds, additives or diluents medicinally approved. The single- and double-stranded, linear DNAs of this invention may be used as an injection either in the form of an aqueous solution as described above or in the form of solid obtained by lyophylizing the same.

The single- and double-stranded, linear DNAs of this invention, when orally administered to animal and 25 human bodies, may be used in the form of capsules, granules, pills, fine granules, tablets or syrup, as in the

The fact that a specific base sequence in a DNA molecule has an important effect upon its case of common drugs. immunostimulatory activity has not been known at all, and this is a completely new finding.

DNAs of this invention enhance the production of interferon and macrophage activating factor, thus activating NK cells and macrophages, also enhance the production of colony-stimulating factor, promote the proliferation of lymphocytes, and are therefore considered to exhibit a wide range of immunostimulatory activity. In addition, these DNAs proved to be very efficacious remedies against experimental tumors, and experimental models for immunodeficiency diseases and for autoimmune diseases, through their immunostimulatory activity. Furthermore, the acute toxicity of these DNAs is much lower than that of synthetic RNA; thus these DNAs are expected to be highly efficacious and useful remedies against malignant tumors, various auto-immune diseases, immunodeficiency diseases and infectious diseases.

EXAMPLES

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Detailed below are Test Methods and Examples, in which guanine, adenine, cytosine and thymine (bases contained in nucleic acids) are abbreviated as G, A, C and T, respectively, and the nucleotide sequence in each DNA molecule is represented so that the left side is 5'-terminal and the right side is 3'terminal. The base sequence in each DNA used in the Examples is shown in the Sequence List appended at the end of this specification. The complex of polyinosinic acid and polycytidylic acid (hereinafter abbreviated as polyl:C), which is a synthetic RNA used as the control remedy, was purchased from Yamasa Shoyu.

Test Method 1 (In-vitro tests on the augmentation of mouse NK-cell activity, on the production of interferon. and on the production of macrophage activating factor)

The tests were performed according to the known method described in the following literature:

Yamamoto, S., et al.: Jpn. J. Cancer Res., 79, 866-873 (1988) provided that the NK-cell activity was measured by a four-hour 51Cr release assay using YAC-1 cells as targets, and the result was expressed by the average of triplicate measurements and the standard sevial on The effector to target ratio was IOO:I unless otherwise stated.

Test Method 2 (Titration of colony-stimulating factor)

A solution of DNA in PBS was administered to MRL MPJ-lpr mice (female, six weeks old) subcutaneously three times per week, and the amount of protein in the urine was calculated from the urine volume excreted during I6 hours and the protein concentration therein.

Test Method 7 (Acute toxicity test in mice)

DNA or RNA was administered to ICR mice (10 mice per group) intravenously (iv) or intraperitoneally (ip), and the number of living mice was counted 24 hours after administration. Graphs were made in which the ratio of living mice was plotted against the amount of DNA or RNA administered, and the amount of DNA or RNA per body weight that will kill half of the mice was estimated - LD₅₀) mg/kg).

Test Method 8 (Test of efficacy against mice infected with LP-BM5 viruses)

Stock solution of LP-BM5 viruses (0.5 ml) was injected intraperitoneally to each of C57BL/I0 mice (15 five weeks old). From the next day, mice were fed with drinking water ad libitum to which azidothymidine (AZT) used as a control drug, was added at a concentration of 0.5 mg/ml. DNA was administered intraperitoneally every day in an amount of I mg as a solution in PBS. Five weeks after the virus infection, the spleen cells obtained from the mice were suspended in RPMII640 medium containing I0% FCS at a cell concentration of I x I0⁷/ml. After culturing the cells at 37° C for 20 hours under 5% CO₂ in the presence of IL-2 (1000 U/ml, a product of Genzyme Corp.), NK cell activity was measured.

Test Method 9 (Quantitation of DNA)

DNA was dissolved in 0.2mM phosphate buffer (pH 7.0), and the absorbance at 260 nm was measured with the same buffer used as reference. DNA concentration was determined by assuming that the concentration of DNA that gave an absorbance of 1 was 20 μg/ml.

Example 1 (Preparation-of tablets containing, as active ingredient, a single-stranded, linear DNA of this invention)

A mixture of 5 g of sodium salt of a single-stranded, linear DNA (Sequence 1), 53 g of lactose, 50 g of corn starch and 35 g of crystalline cellulose was kneaded with a solution of 5 g hydroxypropylcellulose in 10 ml water to form granules, which were dried at 50°C for four hours. Magnesium stearate (2 g) was then admixed, and the mixture was compressed into tablets (each weighing 200 mg) by the use of a tabletting machine.

Example 2 (Preparation of capsules containing, as active ingredient, a single-stranded, linear DNA of this invention)

A mixture of 5 g of sodium salt of a single-stranded, linear DNA (Sequence 1), 124 g of lactose, 90 g of corn starch, 70 g of crystalline cellulose and 11 g of magnesium stearate was filled into hard gelatin capsules (300 mg in each) by the use of a capsule filling machine, thus giving capsules.

Example 3 (Preparation of parenteral injections containing, as active ingredient, a single-stranded, linear DNA of this invention)

One gram of sodium salt of a single-stranded, linear DNA (Sequence 1) and 0.5 g of sodium chloride were dissolved in 1 liter of distilled water for injection, and the mixture was filtered and sterilized, thus giving injections.

Example 4

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The fact that DNAs containing palindromic structure have stronger immunopharmacological activity than those containing no palindromic structure was demonstrated by the following experiments.

A singl -stranded, linear DNA (Sequence 1) with a base number of 45 containing a palindromic structure (GACGTC) had a strong activity to augm nt the mous NK-cell activity (Table 1). In contrast, a single-stranded, linear DNA (Sequence 2) with a bas numb r of 45 containing no palindromic structure had a weaker activity to augment the mouse NK-cell activity (Tabl 1).

palindromic structure was replaced with a reversed sequence of nucleotides (TG), had a strong activity to augment the NK-cell activity like the DNA (Sequence 3) (Table 3).

The DNA (Sequence 7), which is of much the same structure as the DNA (Sequence 3) except that one nucleotide (C) in the palindromic structure is lacking, had a markedly weaker activity to augment the NK-cell activity than the DNA (Sequence 3) (Table 3). However, the DNA (Sequence 8), which is of much the same structure as the DNA (Sequence 3) except that one nucleotide (C) in the portion other than the palindromic structure is lacking had a strong activity to augment the NK-cell activity like the DNA (

The DNA (Sequence 9), which is of much the same structure as the DNA (Sequence 3) except that Sequence 3) (Table 3). the palindromic structure is translocated to the 5'-terminal, had as strong an activity to augment the NK-cell activity as that of the DNA (Sequence 3) (Table 3); however, the DNA (Sequence 10), in which the palindromic structure is translocated to the central position, had a stronger activity to augment the NK-cell activity than that of the DNA (Sequence 3) (Table 4).

Table 3

15	Table 3				
	Test Sample	Number of Bases	NK-Cell Activity		
			14.1±2.0		
20	Control		43.8±2.6		
	Sequence 3	30			
	Sequence 5	30	15.8±1.5		
25	Sequence 6	30	45.3±2.3		
	Sequence 7	29	15.4±1.4		
		. 29	50.9±1.9		
30	Sequence 8		110 to 3		

Each DNA was added to spleen cells to a final concentration of 50 µg/ml.

Table 4

	Test Sample	Number of Bases	NK-Cell Activity
40		Ŋ.	14.2±0.9
	Control		
	Sequence 3	30	43.4±2.1
		20:	45.8±2.6
45	Sequence 9	30	
•	Sequence 10	30	51.4±2.8
		1	- colle to a

Each DNA was added to spleen cells to a final concentration of 50 µg/ml.

It was d monstrated from the abov results that the presence of the palindromic structure (GACGTC) 55 has an important effect upon the DNA's activity to augment th NK-cell activity, and that a higher activity can be obtain d if the palindromic structure is located in the central position of the DNA chain.

Example 5

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Example 6

The following experiment demonstrated that DNAs containing a palindromic structure with a base number of 8 or 10 also have a high immunopharmacological activity. The DNA (Sequence 21) containing a 5 palindromic structure (GACGTC), and the DNAs (Sequences 22 and 23) containing an expanded palindromic structure (GGACGTCC, CGGACGTCCG), had a strong activity to augment the NK-cell activity (Table 7). However, the DNA (Sequence 24) containing a curtailed GACGTC structure (ACGT) had only a low activity.

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Table 7

Test Sample Palindrome NK-Cell Activity Control 14.9±1.0 Sequence 24 ACGT 15.0±0.3 Sequence 21 GACGTC 42.3±1.1 Sequence 22 GGACGTCC 45.5±2.2			
Control 14.9±1.0 Sequence 24 ACGT 15.0±0.3 Sequence 21 GACGTC 42.3±1.1 Sequence 22 GGACGTCC 45.5±2.2 Sequence 23 GGACGTCC 53.6±1.5	Test Sample	Palindrome	NK-Cell Activity
Sequence 24 ACGT 15.0±0.3 Sequence 21 GACGTC 42.3±1.1 Sequence 22 GGACGTCC 45.5±2.2 Sequence 23 GGACGTCC 53.6±1.5			14.9±1.0
Sequence 21 GACGTC 42.3±1.1 Sequence 22 GGACGTCC 45.5±2.2 52.6±1.5 53.6±1.5		ACGT	15.0±0.3
Sequence 22 GGACGTCC 45.5±2.2		GACGTC	42.3±1.1
52 6+1.5		GGACGTCC	45.5±2.2
	Sequence 23	CGGACGTCCG	52.6±1.5

final concentration of 50 μ g/ml.

Example 7

The fact that the immunopharmacological activity of DNAs containing a palindromic structure depends on the molecular length was discovered from the results of the following experiments.

Comparison of the activity to augment the NK-cell activity among DNAs (Sequences 25 through 32) with a palindromic structure in the central position and having a variety of molecular lengths (number of bases: 6 to 80) showed that the activity was observed only in the DNAs having 10 or more bases, increased with the number of bases, and changed little when the number of bases exceeded 45 (Table 8).

activity to augment the NK-cell activity (Table 10).

Table 10

Test Sample	Palindrome	NK-Cell Activity	
		14.2±0.5	
Control	GACGTC	45.1±3.2	
Sequence 37	CACGTG	47.6±2.9	
Sequence 38	AACGTT	42.5±3.2	
Sequence 39	AACGII	11- +0 3	

Each DNA was added to spleen cells to a final concentration of 20 µg/ml.

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Example 9

The palindrome-containing DNA which carries only guanine and cytosine as component base (Sequence 40) and the one which carries only adenine and thymine as component base (Sequence 41) had only a weak activity to augment the NK-cell activity (Table 11).

Table 11

			·	
	Comple	Palindrome	Component Base	NK-Cell Activity
30	Test Sample			14.4±1.6
	Control			46.3±2.5
	Sequence 3	GACGTC	G, A, C, T	
35	Sequence 40	GCGCGC	G, C	14.9±1.3
		ATATAT	A, T.	14.5±1.4
	Sequence 41			lls to a

Each DNA was added to spleen cells to a final concentration of 50 μ g/ml.

It was demonstrated from the above results that, in order for a DNA to exhibit a satisfactory immunopharmacological activity, it must contain at least one palindrome composed of six or more bases; the total number of bases contained therein must be ten or more; and the sequences entirely composed of repetition of GC- or AT-array are unfavorable.

Example 10 50

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Single-stranded, linear DNAs containing palindromic structure induced interferon (hereinafter abbreviated as IFN) and macrophage activating factor (hereinafter abbreviated as MAF).

The DNA (Sequence 1) containing a palindromic structure with a base number of 45 had an in-vitro activity to induce IFN and MAF from mouse spl en cells, but the activity of the DNA (Sequence 2) without palindromic structur was weaker (Table 12).

Example 12

ConA-stimulated proliferation of spleen cells was promoted in mice to which 5 mg of the palindromecontaining DNA (Sequence 1) with a base number of 45 had been administered (Table I4). However, no such activity was observed with the DNA (Sequence 2) without palindromic structure.

Table 14

	Test Sample	Days after	Administration	S.I.
10				42.3
	Control	•		76.1**
	Sequence 1		1	
15			2	66.7*
			3	47.9#
			1	45.5
20	Sequence 2		2	43.1
			3 .	42.0
25	*: p<0.05,	**: 9<0.01	(by Student's	t-test)

The above results demonstrated that DNAs containing palindromic structure exhibit a variety of 30 immnopharmacological activities.

Example 13

When the palindrome-containing DNA (Sequence 1) with a base number of 45 was administered to mice bearing IMC carcinoma tumors, dose-dependent suppression of the tumor weight (namely, antitumor activity) was observed (Table I5). However, the DNA (Sequence 2) with a base number of 45 without palindromic structure had only a weak antitumor activity (Table 15).

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Example 15

Administration of the single-stranded, linear DNA (Sequence 1) containing a palindromic structure to the autoimmune disease model, MRL:MPJ-lpr mice which suffer spontaneous outbreak of such diseases, suppressed the amount of protein excreted in the urine (Table 17). However, no such activity was observed with the DNA (Sequence 2) without palindromic structure.

Table 17

*.		14510	
	Dose	Protein Content	in the Urine (mg)
Test Sample	(mg/kg)	At the Start	After 4 Neeks
Control	0	0.7±0.5	4.0±2.6
Sequence 1	0.03	0.7±0.5	3.1±1.8
3eque.ioc	0.3	0.7±0.5	3.3±1.S
	3	0.6±0.4	3.5±1.3
Sequence 2	0.03	0.7±0.6	4.1±2.3
Sequence 2	0.3	0.5±0.4	3.5±1.6
	3	0.7±0.5	3.9±3.0

Table 17 (contd.)

		Protein Content	in the Urine (mg)
Test Sample	Dose (mg/kg)	After 8 Weeks	After 12 Weeks
Control	0	7.1±13.5	13.6±22.2
Sequence 1	0.03	2.5±3.1	4.5±6.1
Sequence .	0.3	1.7±1.1*	8.6±12.3
	3	3.2±5.8	9.7±16.1
Sequence 2	0.03	7.3±9.5	12.6±6.2
Sedaence -	0.3	5.8±5.2	10.7±12.3
•	3	5.4±4.0	13.0±18.3

^{*:} p<0.05 (Student's t-tgest)

The abov results demonstrated that synthetic DNAs containing palindromic structure have not only immunopharmacological activities but also therapeutic effects on various diseases which are known to be susceptible to drugs with immunopharmacological activity.

Table 20

	Test Sample [alindrome	NK-Call Activity
	Control		12.3±0.6
•	Sequence 11	AACGTT	50.0±1.8
	Sequence 12	AGCGCT	43.2=1.9
•		CGATCG	47.7±1.4
	Sequence 49	ATCGAT	47.0±1.6
	Sequence 50	TCGCGA	47.0±2.0
	Sequence 51	GCGCGC	46.9±1.2
	Sequence 52	CGTACG	46.5±1.9
	Sequence 53	AGCGCT	45.7±1.0
	Sequence 54	CGGCCG	44.2±1.7
	Sequence 55	GACGTC	42.1±1.5
	Sequence 3	GTCGAC	42.0±1.3
	Sequence 56		40.1±1.4
	Sequence 57	CGCGCG	39.5±1.5
	Sequence 58	ACGCGT	20.1±0.8
	Sequence 59		19.9±0.5
	Sequence 50		18.3±0.4
•	Sequence 61		18.0±0.7
	Sequence 62		17.7±0.6
	Sequence 6		5.0 5
	Sequence 6		
	Sequence 6		47.0.0.7
5	Sequence 6	6 CCTAGG	

Each DNA was added to spleen cells to a final concentration of 50 µg/ml.

Example 19

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In mice infected with LP-BM5 viruses (model animals for immunodeficiency diseases), the NK-cell ss activity of the lymphocyt s was not enhanced by IL-2 stimulation unlike in normal mice; however, in the virus-infected mic to which the single-stranded, linear DNA (Sequenc 1) containing a palindromic structure had been administered, IL-2 was able to enhance the NK-cell activity of the lymphocytes (Table 21). The degree of enhancement was nearly the same as that observ d in the IL-2-stimulated lymphocytes

Table 22

			NK-Cell Activity	IFM Titer (U/ml)	
	House	Test Sample		< 4	
5	BALB/c	Control	5.2±0.5	4.00	
	5,	Sequence 1	15.4±0.7	120	
			5.4±0.3	٠4.	
10		Sequence 2		16	
	SCID	Control	18.3±1.0	256	
	•	Sequence 1	43.9±1.5	250	
		*	20.0±1.1	1.6	
15		Sequence 2	spleen cells	to a final	

Each DNA was added to spleen cells to a final concentration of 20 µg/ml.

NK-cell activity was measured at an E:T ratio of 25:1.

It was demonstrated from the results obtained in Examples 19 and 20 that the single-stranded, linear DNAs of this invention containing palindromic structure are capable of restoring (at least partially) the 25 immunological functions of immunodeficiency disease model mice.

Example 21

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Acute toxicity of the synthetic DNAs (Sequences 1 and 3) with a base number of 45 and 30, respectively, was remarkably low as compared to that of the synthetic RNA (polyl:C) used as control (Table 23).

Table 23

Table 23	
Administration Route	LD ₅₀ (mg/kg)
	>500
)	>1.000
ip	
iv	>500
	>1000
	8
ip	30
	Administration Route iv ip iv iv iv

In addition, intrap ritoneal administration of each of the synthetic DNAs (Sequences 10, 11, 12, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 42, 43, 44, 45, 49, 55 50, 51, 52, 53, 54, 55, 56, 57 and 58) to DDY mice (each group consisting of ten heads), caused no death or body weight loss in any of the mice tested in one-week obs rvation period.

Sequence No.: 4

Length of sequence: 30

Type of sequence: Nucleic acid:

Type of Chain : Single-stranded

Topologty: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence:

GGTGACGGCA CCACGACGGC CACCGTGCTG

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Sequence No.: 5

Length of sequence: 30

Type of sequence: Nucleic acid

Type of Chain: Single-stranded

Topologty: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence:

35 ACCGATGACT GCGCCGGTGA CGGCACCACG

Sequence No.: 6

Length of sequence: 30

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

Topologty: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (GACGTC)

ACCGATGACG TCGCCGTGGA CGGCACCACG

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Sequence No.: 10

Length of sequence: 30

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

Topologty: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (GACGTC)

ACCACGACCG ATGACGTCGC CGGTGACGGC

Sequence No.: 11

Length of sequence: 30

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

Topologty: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (AACGTT)

ACCGATAACG TTGCCGGTGA CGGCACCACG

Sequence No.: 12

Length of sequence: 30

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

Topologty: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (AGCGCT)

ACCGATAGCG CTGCCGGTGA CGGCACCACG

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Sequence No.: 16

Length of sequence: 30

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

Topologty: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (ATCGAT)

TCGGTGATCG ATATGTCGCA GGACCCGGTC

Sequence No.: 17

Length of sequence: 30

Type of sequence: Nucleic acid

25 Type of Chain : Single-stranded

Topologty: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (TCGCGA)

TCGGTGTCGC GAATGTCGCA GGACCCGGTC

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Sequence No.: 18

Length of sequence: 30

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

45 Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (GCGCGC)

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AAAAGAAGTG GGGACGTCTT ACGATCACCA

Sequence No.: 22

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Length of sequence: 30

Type of sequence: Nucleic acid 10

Type of Chain : Single-stranded

Topology: Linear 15

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (GGACGTCC)

AAAAGAAGTG GGGACGTCCT ACGATCACCA

Sequence No.: 23 25

Length of sequence: 30

Type of sequence: Nucleic acid

Type of Chain : Single-stranded 30

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (CGGACGTCCG)

AAAAGAAGTG CGGACGTCCG ACGATCACCA

Sequence No.: 24

Length of sequence: 30

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (ACGT)

CCGATGACGT CGCCG

i

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Sequence No.: 28

Length of sequence: 20

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (GACGTC)

GACCGATGAC GTCGCCGGTG

Sequence No.: 29

Length of sequence: 30

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (GACGTC)

TGACAGACCG ATGACGTCGC CGGTGGACGG

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Sequence No.: 30

Length of sequence: 45

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

50 Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (GACGTC)

Features of sequence: P: Contains palindrome (GACGTC) ACCGATGACG TCGCGACGTC CGGCACCACG ACGGCCACCG TGCTG

Sequence No.: 34

Length of sequence: 45 10

Type of sequence: Nucleic acid

Type of Chain : Single-stranded 15

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (GACGTC)

ACCGATGACG TCGCGACGTC CGGACGTCCG ACGGCCACCG TGCTG

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Sequence No.: 35

Length of sequence: 45

Type of sequence: Nucleic acid 30

Type of Chain : Single-stranded

Topology: Linear 35

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (GACGTC)

ACCGATGACG TCGCGACGTC CGGACGTCCG GACGTCACCG TGCTG

Sequence No.: 36

Length of sequence: 45

Type of sequence: Nucleic acid

Type of Chain : Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA 55

Features of sequence: P: Contains palindrome (AACGTT) AACGTTAACG TTAACGTTAA CGTTAACGTT

Sequence No.: 40

Length of sequence: 30 10

Type of sequence: Nucleic acid

Type of chain: Single-stranded 15

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (GCGCGC)

Sequence No.: 41

Length of sequence: 30

Type of sequence: Nucleic acid

Type of chain: Single-stranded

Topology: Linear 35

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (ATATAT)

ATATATATAT ATATATATAT ATATATATAT

Sequence No.: 42

Length of sequence: 45

Type of sequence: Nucleic acid

Type of chain: Double- stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (GACGTC) TTTTTTTTT TTGACGTCTT TTTTTTTTT 10 Sequence No.: 46 Length of sequence: 30 Type of sequence: Nucleic acid Type of chain: Single-stranded Topology: Linear Kind of sequence: Other nucleic acid; synthetic DNA 20 Features of sequence: P: Contains palindrome (GACGTC) CCCCCCCC CCGACGTCCC CCCCCCCCC 25 Sequence No.: 47 Length of sequence: 30 Type of sequence: Nucleic acid Type of chain: Single-stranded Topology: Linear Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome(GACGTC) 40 GCGCGCGCG GCGACGTCGC GCGCGCGCGC Sequence No.: 48 Length of sequence: 30 Type of sequence: Nucleic acid Type of chain: Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (TCGCGA) ACCGATTCGC GAGCCGGTGA CGGCACCACG

Sequence No.: 52

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Length of sequence: 30

Type of sequence: Nucleic acid 15 Type of chain: Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (GCGCGC)

ACCGATGCGC GCGCCGGTGA CGGCACCACG

Sequence No.: 53

Length of sequence: 30

Type of sequence: Nucleic acid

Type of chain: Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA

Features of sequence: P: Contains palindrome (CGTACG) 40

ACCGATCGTA CGGCCGGTGA .CGGCACCACG

Sequence No.: 54

Length of sequence: 30

Type of sequence: Nucleic acid

Type of chain: Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (CGCGCG) ACCGATCGCG CGGCCGGTGA CGGCACCACG

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Sequence No.: 58

Length of sequence: 30

Type of sequence: Nucleic acid 15

Type of chain: Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (ACGCGT)

ACCGATACGC GTGCCGGTGA CGGCACCACG 25

Sequence No.: 59

30 Length of sequence: 30

Type of sequence: Nucleic acid

Type of chain: Single-stranded 35

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (AAGCTT)

ACCGATAAGC TTGCCGGTGA CGGCACCACG

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Sequence No.: 60

Length of sequence: 30

Type of sequence: Nucleic acid 50

Type of chain: Single-stranded

Topology: Linear 55

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (GAATTC) ACCGATGAAT TCGCCGGTGA CGGCACCACG

Sequence No.: 64

Length of sequence: 30

Type of sequence: Nucleic acid

Type of chain: Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (CTGCAG) 20 ACCGATCTGC AGGCCGGTGA CGGCACCACG

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Sequence No.: 65

Length of sequence: 30

Type of sequence: Nucleic acid Type of chain: Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA Features of sequence: P: Contains palindrome (AAATTT)

ACCGATAAAT TTGCCGGTGA CGGCACCACG 40

Sequence No.: 66

Length of sequence: 30

Type of sequence: Nucleic acid

Type of chain: Single-stranded

Topology: Linear

Kind of sequence: Other nucleic acid; synthetic DNA 55

- general formula (I) is 5'-TCGCGA-3' (wherein G is deoxyguanylic acid, A is deoxyadenylic acid, C is deoxycytidylic acid, and T is deoxythymidylic acid).
- 12. An immunostimulatory remedy as described in Claim 1 or 2, wherein the structure represented by the general formula (I) is 5'-AACGTT-3' (wherein G is deoxyguanylic acid, A is deoxyadenylic acid, C is deoxycytidylic acid, and T is deoxythymidylic acid).
 - 13. An immunostimulatory remedy as described in Claim 1 or 2, wherein the structure represented by the general formula (I) is 5'-GCGCGC-3' (wherein G is deoxyguanylic acid, and C is deoxycytidylic acid).
- 14. An immunostimulatory remedy as described in Claim 1 or 2, wherein the structure represented by the general formula (I) is 5'-CGTACG-3' (wherein G is deoxyguanylic acid, A is deoxyadenylic acid, C is deoxycytidylic acid, and T is deoxythymidylic acid).
- 15. An immunostimulatory remedy as described in Claim 1 or 2, wherein the structure represented by the general formula (I) is 5'-CGGCCG-3' (wherein G is deoxyguanylic acid, and C is deoxycytidylic acid).
- 16. An immunostimulatory remedy as described in Claim 1 or 2, wherein the structure represented by the general formula (I) is 5'-GTCGAC-3' (wherein G is deoxyguanylic acid, A is deoxyadenylic acid, C is deoxycytidylic acid, and T is deoxythymidylic acid).
 - 17. An immunostimulatory remedy as described in Claim 1 or 2, wherein the structure represented by the general formula (I) is 5'-CGCGCG-3' (wherein G is deoxyguanylic acid, and C is deoxycytidylic acid).
- 25 18. An immunostimulatory remedy as described in any of Claims 1 through 17, wherein the portion except the structure represented by the general formula (I) is the repeated structure of deoxyguanylic acid.
 - 19. An immunostimulatory remedy as described in Claim 1 or 2, wherein the structure represented by the general formula (I) contains at least one structure of 5'-CG-3' (wherein G is deoxyguanylic acid, and C is deoxycytidylic acid).
 - 20. An immunostimulatory remedy as described in Claim 19, wherein the portion except the structure represented by the generagl formula (I) is the repeated structure of deoxyguanylic acid.
 - 21. An immunostimulatory remedy as described in any one of Claims 1 to 20, for the relief of malignant tumors, infectious dieases, immunodeficiency diseases and autoimmune diseases.

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-			1-2	A61K31/70	
	EP-A-0 300 687 (CITY OF HOPE)				
	* page 2, line 48 - line 49 *		_	1	
Ì	EP-A-0 302 758 (NEW ENGLAND MEDIC	AL CENTER	1-2		
	HOSPITALS)				
	* claims 2-4 *	٠.			
	DE-A-3 744 785 (THEURER K.)		1-2		
	" the whole document "				
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	WORLD PATENTS INDEX LATEST		1-21		
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	Week 9026, Derwent Publications Ltd., Londo	on, GB;			
	AN 90-196689 & JP-A-2 128 691 (NIPPON OIL SEA	L) 17 May 1990			
	* abstract *		1		
			1-21		
	MEDLINE ABSTRACT Number: 903380	13	1-21		
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